Abstract The relationship between the sacoglossan Elysia timida and the ornate wrasse Thalassoma pavo was studied in two laboratory experiments using artificial models. A feeding-preference experiment was conducted to determine whether mollusc extract deterred feeding by T. pavo, by using a "realistic" mollusc model (imitating the colour pattern of E. timida) coated with mollusc extract, and a reference model (without imitation or extract), and fishes collected from two locations. It was observed that fish approached, attacked and ate more reference models than mollusc models. A second feeding-preference experiment was designed with four different coloured models: "realistic" (W), green (G), red (R) and chequered (S) pattern. Both fish populations clearly rejected the S model, and differed in their colour preferences. Although both populations seemed to prefer the R and G models equally, the W model was clearly rejected by the fish that coexist with the mollusc at one site (Mazarrón), but was not rejected by the other population of fish which does not coexist with it (Cabo de Palos). Mazarrón fishes would identify the W model with the presence of a toxic compound during their coexistence, and therefore avoid attacking conspicuously coloured E. timida models as a response to their visual signals. Therefore, it was concluded that extract of E. timida is a deterrent for T. pavo, and its effect is sufficiently noxious that the fish tend to avoid it, so that the ability of the fishes to learn to recognise colours and identify certain colour patterns associated with obnoxious prey provides the molluscs the opportunity to survive by exhibiting a conspicuous coloration.

Introduction

The predator–prey relationship is one of the strongest selection factors in the evolution of behaviour (Csányi and Dóka 1993). Colour-pattern strategy (aposematism or crypsis), chemical defence and shelter background are prey adaptations to avoid predator pressure, while improved visual learning of specific patterns and detoxification ability are predator adaptations to counteract prey abilities (Krebs and Davies 1993). The visibility of an organism’s colour pattern is affected not only by the inherent properties of the pattern, but also by predator vision, hunting tactics, prey behaviour, and the background colour patterns. The effects of these factors are not independent and may vary from place to place (Endler 1978). Bright coloration is often associated with distastefulness, since a conspicuous prey is easily recognised. Predators learn to avoid warning colours effortlessly, but having learned to avoid a prey type they are probably less likely to attack it by mistake if it is conspicuous (Guilford 1985).

It is frequently accepted that many marine molluscs use distasteful molecules to avoid attack by predator fishes. The opisthobranchs are shell-less molluscs with limited mobility and a broad variety of conspicuous coloration. During their evolution they have lost the mechanical protection given by the shell, but they have developed defensive strategies based on the use of chemical substances (Ros 1977; Rosenberg 1991). The principal source of chemical substances is their food; however, some opisthobranchs are able to modify chemical substances that have been sequestered from
their prey or to biosynthesise them de novo (Cimino and Sodano 1993; Fontana et al. 1994).

The sacoglossan *Elysia timida* (Risso 1818) is a white Mediterranean opisthobranch that lives in well lit shallow waters characterised by low-energy hydrodynamic conditions. They are found in areas covered either by stones or a thick layer of sand. Such environments are also typical of the chlorophycean seaweed *Acetabularia acetabulum* (L.) (Marín and Ros 1992; Giménez-Casaldueiro 1999). This mollusc is able to biosynthesise de novo polypropionate, presumably as a chemical defence mechanism (Gavagnin et al. 1994a, b; Cimino and Ghiselin 1998). The ornate wrasse *Thalassoma pavo* L. is a labrid fish very common in south-western Mediterranean coastal regions (García-Charton and Pérez-Ruzafa 2001). It lives in the typical sacoglossan habitat, and feeds on small crustaceans and molluscs (Whitehead et al. 1986), and is therefore a potential predator of *E. timida*. It has been found that the different compounds isolated from this mollusc are toxic for the fish *Gambusia affinis* (commonly used for ecotoxicity tests), but it has not been demonstrated that the polypropionate deter predator fish that cohabit with the mollusc (see Gavagnin et al. 1994b).

The main objective of this paper is to demonstrate the capacity of *E. timida* to deter the marine fish *T. pavo*, and the effectiveness of the mollusc coloration as a warning signal. The mollusc extract and the acceptance of four coloured models (three of them based on the natural mollusc colour and design; and the last one showing an “unrealistic” pattern) were tested for fish populations coming from two different locations in south-eastern Spain.

### Materials and methods

#### Animal collection and fish-holding conditions

Using hand nets, the wrasses were collected from two localities on the coast of Murcia (south-eastern Spain) (Fig. 1): (1) at Cabo de Palos (37°38’N, 0°41’W), where the predominantly rocky bottom is formed by boulders of different sizes with clumps of the seagrass *Posidonia oceanica* interspersed in the rocky matrix; the sacoglossan is rare in this area, probably owing to the heavy wave action prevailing in this zone; (2) Mazarroñ Bay (37°33’N, 1°16’W), a coastal area protected from wave exposure where the sacoglossan and the seaweed *Acetabularia acetabulum* are particularly common and abundant (Marín and Ros 1992). The two areas are separated by 40 km.

The fish were immediately transported to the laboratory, where they were placed in a tank containing a silica–sand substratum and flow-through well water (20°C, salinity 36‰), and exposed to a 12-h light:12-h dark illumination regime. While in captivity, the fish were fed ad libitum two times daily with the same artificial food as that used in the experiments (described below).

The sacoglossan specimens were collected from Mazarroñ Bay at depths of 1–3 m, and were transferred to the laboratory in an icebox within 3 h.

#### Artificial models

The artificial sacoglossan models were based on previous experiments and were similar to those described by Magurran and Guirling (1986), Dugatkin and Godin (1992) and Giménez-Casaldueiro et al. (1999). Each model consisted of two parts: the main body and the outer skin. The main body was made from triturated sardine with 4% of carrageenan (Sigma 1013) (Fig. 2A, a). The ingredients were mixed by stirring and then heated in a microwave oven for 60 s. The outer skin (Fig. 2A, b) was composed of two pieces of printed colour paper 0.7 x 1.5 cm in size, created with a computer design programme. Five different designs were placed on the artificial diet: realistic (W), coloured (R and G), unrealistic (S) and control (X) models. W models were based on the natural physical characteristics of *E. timida* (mollusc colour and design; Fig. 2A, W). In addition to this realistic model, two more models were made with the sacoglossan pattern but with different colours: the R model had red coloration (Fig. 2A, R), and the G model had green coloration (Fig. 2A, G). The “unrealistic” S model had a black-and-white chequered pattern (Fig. 2A, S). Finally, the X model had the same main body and size characteristics as previously, but the outer skin had no design on it, this model being used as a control in the chemical feeding-avoidance experiment (see below) (Fig. 2A, X).

To test the deterrent ability of *E. timida*, mollusc extract was used to coat the W models. Approximately 300 sacoglossans were collected in Mazarroñ Bay and subsequently extracted in acetone, and the solvent evaporated under vacuum. The amount of extract used for each model was estimated according to Gavagnin et al. (1994a). The extract concentration was calculated as milligrams per gram dry weight of animal tissue, using an empirically derived equation for animal weight: $DW = 8.55WW + 4.55$ ($r = 0.82$), where $DW$ is the dry weight and $WW$ is the wet weight (both

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**Fig. 1** Fish-sampling sites in the south-western Mediterranean Sea. *Asterisk* Mazarroñ; *filled circle* Cabo de Palos
Experimental apparatus and video monitoring

The experiments were conducted using a well illuminated tank (110×62×40 cm). The experimental tank, based on that used by Godin and Crossman (1994), was divided into two compartments (Fig. 2B). One compartment contained the fishes, and the other was equipped with hangers upon which the different artificial mollusc models could be placed. The compartments were separated from each other by a removable opaque partition, so that the fishes were not able to see the models prior to the start of the experiment. This opaque partition was raised prior to each trial, to allow the fish to enter the compartment containing the models. The tank was covered on three sides by light blue cardboard, and a video camera was installed on the fourth side. Videotapes were recorded continuously for 20 s after the removal of the partition. Tapes were analysed later by means of a video recorder.

Chemical feeding-avoidance experiment

To determine whether mollusc extract deterred feeding by T. pavo, a feeding-preference experiment was performed in the laboratory. W-model and X-model molluscs were used, the former coated with mollusc extract, and the latter with ether. Each trial consisted of an equal number of each model; three W models and three X models were offered to five fishes at the same time during a 20-s period. Experimental trials were completed for each location, for a total of 36 trials. Data collected included the number of approaches, number of effective interactions (number of bites in each approach), and total number of fully-eaten models.

Colour pattern preference experiment

To measure the effectiveness of the colour pattern as a warning signal for fishes, a feeding-preference experiment was designed, based on Ritland (1994). W, R, G, and S models were tested. Eighteen individuals of T. pavo were independently tested in this experiment, nine from each locality. Each specimen of T. pavo was placed in the experimental tank, and two items of each model design were offered simultaneously to the fish during 20 s. The total number of models eaten was scored for each trial.

Data analysis

To test for differences in behaviour between fishes from the two localities, arcsin-transformed data were compared using a t-test (Underwood 1997) for both experiments. To compare the two populations, the measurements of the fishes’ responses (approaches, effective interaction and number of models eaten) were tested independently. Also, the proportion of W models eaten relative to the total number of models eaten [W/(W + G + R + S)] was analysed in the coloration-preference experiment.

Analysing each population independently, every experiment constituted a multi-choice assay, in which the consumption of one type of model was not independent of the consumption of another type of model. The data were analysed by the one-sample Hotelling’s T² test (Roa 1992; Giménez-Casalduero et al. 1999). This multivariate-analysis technique is applicable in the situation in which p food types are offered to the fish in the tank (in the feeding-avoidance experiment, p = 2 and n = 18; in the coloration-preference experiment, p = 4 and n = 9), so that the response variable in such an experiment is a vector X of mean consumptions of dimension p, in which the components (the different models) are correlated. The null hypothesis to be tested for both experiments was that the consumer has no food preference, which is equivalent to testing whether the components of the mean vector X are all equal to a constant k (the overall mean). The statistical test to prove this hypothesis is Hotelling’s statistic \( T^2 = n(X - k)S^{-1}(X - k) \), where S⁻¹ is the inverse of the sampling variance-covariance matrix, and (X – k) is the vector of differences between the sample means and the hypothesised constant. When the null hypothesis is true, \( T^2 \) has the F distribution with p and n–p degrees of freedom. When the null hypothesis of “no preference” is rejected, those food types with means greater than the constant k (no preference) are suspected to be preferred; and the opposite is true for foods with means less than k. Data were centred on zero to avoid entering the constant k from each datum in order to simplify the identification of preferred and rejected food types, so that the null hypothesis of no preference in this case is whether the components of the mean vector are equal to zero. Since no autogenic changes are expected to occur in the absence of consumers during the experiments, controls were not included (Roa 1992; Manly 1993). The use of dependent variables, of quantities (number of approaches, number of bites and number of models eaten) that estimate...
absolute rather than relative consumption precludes use of the alternative analysis proposed by Lockwood (1998).

Results

Chemical feeding avoidance

Behaviour towards W models did not show significant differences between the two fish populations in respect of either the number of approaches (t = 0.3, P > 0.05), the number of effective interactions (t = 0.6, P > 0.05), or the number of models eaten (t = 0.5, P > 0.05) (Fig. 3). The extract used on the W model seemed to cause rejection by the fishes, since fishes coming from both populations rejected the W models equally for all the parameters examined in the present study. In addition, when each population was analysed independently, we observed that the W models were less inspected than the X models by both T. pavo populations, and that the number of approaches to W models was found to be significantly lower. For each approach to a W model, fishes captured at Cabo de Palos made, on average, 1.6 approaches to an X model, and fishes from Mazarrón made, on average, 1.8 approaches to an X model (Cabo de Palos: T² = 0.65, F₂,₁₆ = 5.19, P < 0.05; Mazarrón: T² = 1.75, F₂,₁₆ = 13.98, P < 0.001) (Table 1; Fig. 3). Furthermore, the number of effective interactions (number of bites relative to number of approaches) was lower for the realistic W models than for the X models for each population (Cabo de Palos: T² = 6.38, F₂,₁₆ = 51.00, P < 0.001; Mazarrón: T² = 6.40, F₂,₁₆ = 51.16, P < 0.001) (Table 1; Fig. 3). Fishes were rather efficient with respect to X models, since 80–90% of approaches resulted in bites, this effectiveness being lower for W models (55% for the Cabo de Palos population and 49% for the Mazarrón population). The number of models eaten at the end of the experiment was similar to that seen for effective attacks, since fishes coming from both populations ate 41% more X models than W models impregnated with toxin (Cabo de Palos: T² = 1.36, F₂,₁₆ = 10.88, P < 0.001; Mazarrón: T² = 1.40, F₂,₁₆ = 11.16, P < 0.001) (Table 1; Fig. 3).

Coloration preferences

Fish behaviour towards W models was compared between the two populations using the number of W models eaten relative to the total number of models eaten each time. The results showed that the population cohabiting with the mollusc, at Mazarrón, rejected the W model, whereas the population from Cabo de Palos did not (Fig. 4; Student’s t = 2.7, P < 0.05).

Both fish populations clearly rejected the chequerboard design of the S model: the Mazarrón population ate only 11% of these models, and the Cabo de Palos fishes, only 28% (Table 2; Fig. 4). In addition, fishes from Cabo de Palos and from Mazarrón differed as to their colour preferences. Both populations preferred the R and G models equally, but the W model was clearly rejected by only the Mazarrón fishes (Table 2). The Mazarrón population ate the R and G models with the same intensity (94% of models eaten), but the proportion of W models eaten was found to be 39%, an intermediate value between the S model in one part, and the R and G models in the other part (T² = 34.66, F₄,₅ = 43.33, P < 0.001). The Cabo de Palos fishes consumed the W model with the same intensity as they did the R and G models (T² = 17.11, F₄,₅ = 21.39, P < 0.01) (Table 2; Fig. 4).

Discussion

The identification of external signals by fish through chemoreceptors can be achieved by odour, taste or chemoreceptive epidermal cells (Baatrup and Doving 1985; Peters et al. 1987, 1990, 1991; Kotrschal 1991). The capacity to recognise odour in the aquatic environment...
Table 1 Results of Hotelling’s $T^2$ test on number of fish approaches, of effective attacks and of models eaten in feeding-preference experiments on Thalassoma pavo from Mazarrón and from Cabo de Palos in respect of the colour-pattern and chemical feeding preference experiments; $W$ + toxin: Elysia timida model impregnated with sacoglossan extract; $X$ + ether: no-design model impregnated with ether. Centred mean is the vector of differences between the sample means and the hypothesised constant overall mean, after centring on zero

<table>
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<th>Population</th>
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<th>df</th>
<th>F</th>
<th>P</th>
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<td>df</td>
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<td></td>
<td>X + ether</td>
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<td>df</td>
<td></td>
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<td></td>
<td>X + ether</td>
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<td>–0.248</td>
<td>0.248</td>
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<td></td>
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<td>X + ether</td>
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<td>1.40</td>
<td>11.16</td>
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<td>Cabo de Palos</td>
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<td>–0.611</td>
<td>0.611</td>
<td>1.36</td>
<td>10.88</td>
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Fig. 4 Mean values (± SE) of percentage of models eaten by T. pavo in both populations. $W$ E. timida model; $R$ red model; $G$ green model; $S$ black-and-white chequered model

has been evaluated for marine and freshwater species (Chivers and Smith 1993; Saglio and Blanc 1993; Valentinić et al. 1994). In accordance with the results of the present study, a rejection of the extract-coated realistic models exists even before tasting by both fish populations. Therefore, it would seem as if a repellent odour were coming from the mollusc extract. The differences between the W and the X models increased when the effective attack of both populations was analysed. Fishes bit the W model less than they did the X model: this could be the synergic effect of repellent odour and bad taste of E. timida extract. The tastings, and subsequent rejections, of the W model were 41% higher than those for the X models. As a consequence of these results, it could be concluded that the individuals of E. timida are deterrent preys for T. pavo, and their effect is sufficiently noxious for the fish to tend to avoid them.

Some sacoglossans have adapted to feeding upon chemically defended plants (Marín and Ros 1989; Paul et al. 1982, 1992), presumably because they avoid predation and incidental ingestion by macroherbivores, or because they sequester secondary metabolites from these seaweeds, which then protect them from their own predators (Hay 1991). For its part, E. timida feeds on host plants, for example Acetabularia acetabulum, that do not contain deterrent metabolites, and therefore it must synthesise its own defence. Ireland and Scheuer (1979) discovered the polypropionate photodeoxytridachione and 9,10-deoxytridachione in the Pacific sacoglossan Placobranchus ocellatus, a metabolic product that is also found in E. timida. Furthermore, E. timida contains the polypropionates 15-norphotodeoxytridachione and iso-9,10-deoxytridachione, chemicals that are ejected in the event of an alarm (Gavagnin et al. 1994b), and found exclusively in this Mediterranean mollusc. For another example, Elysia viridis feeds upon Codium vermilara, which is protected by elysione that is synthesised de novo with a high level of incorporation of propionate (Gavagnin et al. 1994a; Cimino et al. 1999). Muniain et al. (1999) mentioned a non-characterised polypropionate from the mantle of the south Atlantic species Elysia patagonica, recently described by Muniain and Ortea (1997). The sacoglossan lives on the green alga Bryopsis plumose, from which it sequesters intact chloroplasts in the vacuoles of its digestive cells. The presence of chemical metabolites only appears to be associated with this mollusc (Muniain et al. 2001). The presence of the same molecules in species from distinct geographical areas suggests an effective defensive role of these polypropionates against different potential predators in several marine ecosystems, but it has only been demonstrated in the laboratory in a few cases. Generalist herbivores feed on a wide variety of plants and so become more widely dispersed in space. The selective pressure exerted by natural predators might have resulted in the evolution of de novo defences (Hay 1991; Cimino et al. 1999).

The results obtained in the test of coloration preferences seem to indicate that the wrasse T. pavo, regardless of its origin, can discriminate mollusc colour and reject a disruptive colour pattern (model S). The disruptive color pattern consists of a design that breaks the visual contour of the animal, causing the animal to integrate visually into its background elements, but without formal
provides four not mutually exclusive hypotheses to
indistinguishable; Marı´n and Ros 1989). Guilford (1990)
mimetica strategy is common in elysiids (for example,
distinguish the prey from its background; the latter
pattern of a prey ensures that a predator can not easily
as opposed to cryptic coloration, in which the colour
advertise noxious properties to potential predators,
represented a response to the W model similar to that for
G and R models. Fishes captured in Mazarro´n did not
associate distastefulness with red and green colour, but
they did with the realistic white colour pattern.
Bearing in mind that the only difference between both
fish populations was the habitat and the presence or
not of sacoglossans in that habitat, the observed
response of the Mazarrón T. pavo population is likely
because these fishes have learned to avoid the E. timida
colour pattern in their own habitat, by discriminating
mollusc colours and associating the E. timida colour
with noxuousness. The E. timida design seems to be a
particularly optimum aposematic coloration design to
avoid the fish attack. These results are in agreement
with those of Giménez-Casalduevo et al. (1999), who
indicated that the common colours of nudibranchs are
recognisable by fishes and can be associated with
noxious prey. The observed differences in the beha-
viour of the Cabo de Palos fishes between the two
experiments with respect to the W model could be
attributed to the presence of the extract used in the
feeding-avoidance experiment, instead of coloration or
design. We cannot conclude that T. pavo can dis-
criminate between green and red wavelengths.

Aposematic coloration is used by prey species to
advertise noxious properties to potential predators, as
opposed to cryptic coloration, in which the colour
pattern of a prey ensures that a predator can not easily
distinguish the prey from its background; the latter
strategy is common in elysids (for example, Bosellia
mimetica, which feeds on Halimeda tuna and is truly
indistinguishable; Marin and Ros 1989). Guilford (1990)
provides four not mutually exclusive hypotheses to
explain why organisms use conspicuousness for warning:
(1) predators learn to associate distastefulness with
conspicious colour pattern more rapidly than with a
cryptic colour pattern; (2) certain specific colour pat-
tterns are easier to associate with distastefulness than
others; (3) new patterns are easier to learn; and (4)
conspicious coloration will allow fewer recognition
errors than a cryptic one. Also, Edmunds (1991) gives four
criteria by which to demonstrate that a mollusc is
aposematic for a potential predator, if: (1) it is suffi-
ciently noxious that the predator will not eat it; (2) it is
conspicuously coloured; (3) the predators avoid attack-
ing it because of its signals; (4) these conspicuous signals
provide better protection to the individual or to its genes
than other signals would. While many authors have
generally accepted the existence of aposematic colora-
tion in marine molluscs, these features have often been
described only from a human perspective.

In summary, the present study shows experimental
evidence of aposematism by E. timida against its po-
tential predator T. pavo, after demonstrating that the
sacoglossan polypropionates are effective deterrents for
this fish species. Therefore, the ability of fishes to rec-
ognise colours and identify certain colour patterns with
harmful prey allows the molluscs to survive by exhibit-
ing a conspicuous coloration. Additional research is
needed to measure the actual rate of predation avoid-
ance by potential fish predators on E. timida under
natural conditions in the field, thus estimating the rela-
tive importance of mortality due to predation for the
population dynamics of this sacoglossan.

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clare that the experiments complied with the current laws of the
country in which the experiments were performed.

References

Baatrup E, Doving KB (1985) Physiological studies on solitary
receptors of the oral disc papillae in the adult brook lamprey,
Lampetra planeri (Bloch). Chem Senses 10:559–566

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<td>R</td>
<td>G</td>
<td>S</td>
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Table 2 Results of Hotelling's T^2 test on data from feeding-preference experiments on T. pavo from Mazarrón and from Cabo de Palos in respect of the colour pattern preference experiment; W E. timida model; R red model; G green model; S black-and-white chequered model. Centred mean is the vector of differences between the sample means and the hypothesised constant overall mean, after centring on zero.